

# Attribution of direct ozone radiative forcing to spatially resolved emissions

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[1] Quantifying the dependence of ozone direct radiative forcing (DRF) on the mixture and spatial distribution of precursor emissions is a key step towards understanding the impact of air quality standards on climate. We use here a combination of satellite observations of ozone and its radiative effect in conjunction with an adjoint chemical transport model to determine the ozone DRF due to global, anthropogenic NO<sub>x</sub>, CO, and non-methane hydrocarbons (NMHC) emissions regionally at 2° × 2.5° regions resolution. We show that 8% of the ozone DRF from the sum of all these emissions can be attributed to 15 regions, which are predominantly located in China and the United States (US). To achieve an equivalent reduction in ozone DRF, necessary emission reductions for each precursor vary intra-continently by a factor of 3–10 and globally by over an order of magnitude. The contribution of NO<sub>x</sub> emissions to ozone DRF relative to CO and NMHC emissions within individual regions varies globally by nearly a factor of two. **Citation:** Bowman, K., and D. K. Henze (2012), Attribution of direct ozone radiative forcing to spatially resolved emissions, *Geophys. Res. Lett.*, 39, L22704, doi:10.1029/2012GL053274.

## 1. Introduction

[2] Air quality and climate co-benefit strategies for ozone are complicated by the complex spatio-temporal structure of tropospheric ozone and the non-linear chemistry relating precursor emissions to ozone distributions [Forster *et al.*, 2007; Levy *et al.*, 2008; Sitch *et al.*, 2007; Kawase *et al.*, 2011]. Previous studies have focused on climate responses to continental-scale radiative forcing (RF) [Shindell and Faluvegi, 2009], the role of sectorally aggregated [Unger *et al.*, 2010; Fuglestad *et al.*, 2008] and continental-scale [Naik *et al.*, 2005; Berntsen *et al.*, 2006; Stevenson and Derwent, 2009] changes in precursor emissions on RF, and the degree to which increases in CH<sub>4</sub> radiative forcing following NO<sub>x</sub> reductions can offset reduced ozone direct radiative forcing [Naik *et al.*, 2005; Fiore *et al.*, 2008; Derwent *et al.*, 2008; Stevenson and Derwent, 2009].

[3] In this work we address the role of regional (2° × 2.5°) variations in chemical environment and transport in modulating direct ozone radiative forcing (DRF) at intra-continental

scales through a novel approach that uses satellite observations from the Tropospheric Emission Spectrometer (TES) in conjunction with adjoint sensitivity analysis from the GEOS-Chem chemistry and transport model. Observationally constrained radiative forcings are calculated in each grid-cell for more than thirty different emission types, including both natural and sector-specific anthropogenic O<sub>3</sub> precursors; we focus here on anthropogenic NO<sub>x</sub>, CO, and NMHC sources because of their dominant role in ozone photochemistry and air quality (contributions of sector-aggregated NO<sub>x</sub> emissions are provided in Table S1 in Text S1 in the auxiliary material).<sup>1</sup> This level of quantification is made feasible through the use of an adjoint model, which in general is an efficient means of calculating sensitivities with respect to large numbers of model inputs (in this case emissions). This approach, however, only accounts for ozone direct RF, which does not include methane-ozone feedbacks or indirect methane RF. These additional effects have a significant impact on total RF from precursor emissions and would need to be included in any comprehensive air quality-climate co-benefit analysis.

## 2. Methods

### 2.1. Tropospheric O<sub>3</sub> Radiative Effects Using TES IRKs

[4] The Tropospheric Emissions Spectrometer (TES) is a polar sun-synchronous, high resolution (0.1 cm<sup>-1</sup> apodized), infrared Fourier transform spectrometer aboard the NASA Aura satellite with a global repeat cycle of 16 days and an averaged nadir footprint of 5 km × 8 km [Beer, 2006]. Vertical ozone profiles are derived from spectrally-resolved top-of-the-atmosphere (TOA) thermal radiances based on an optimal estimation framework [Bowman *et al.*, 2006]. This relationship between TOA radiances and ozone distributions was first exploited to quantify the greenhouse gas effect of upper tropospheric ozone over clear-sky, oceanic scenes [Worden *et al.*, 2008] but was subsequently formalized for all-sky and land/oceans scenes through the introduction of longwave instantaneous radiative kernels (IRK) defined as

$$\mathbf{k}_i = \frac{\partial F_i}{\partial \mathbf{c}_i} \quad (1)$$

where  $F_i$  is the instantaneous upward TOA flux in atmospheric column location  $i$  integrated across the infrared band in W/m<sup>2</sup>,  $\mathbf{c}_i$  is the TES retrieved ozone profile on  $L$  pressure levels, and  $\mathbf{k}_i$  is the IRK in W/m<sup>2</sup>/ppb of the  $i$ th column.

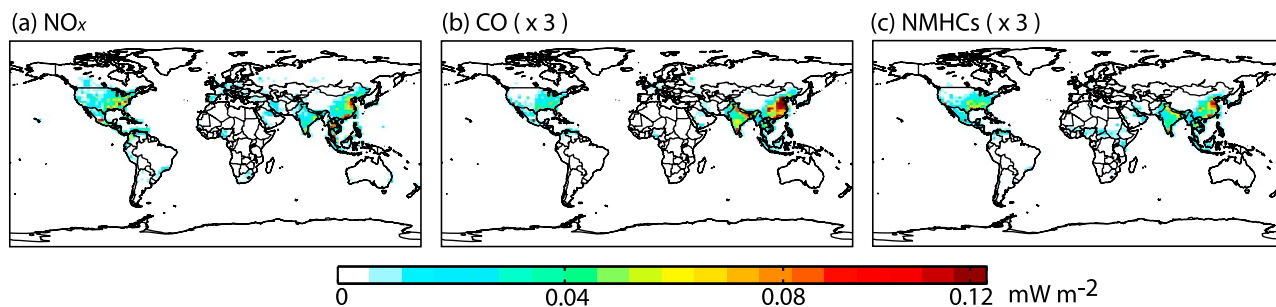
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**Figure 1.** Ozone direct radiative forcing (DRF),  $\lambda$ , as attributed from TES observations for August 2006 emissions to each grid cell of (a) NO<sub>x</sub> (b) CO (scaled by 3) and (c) NMHC (scaled by 3). The color scale is saturated for DRF > 0.12 mW/m<sup>2</sup>.

Under clear-sky scenes, the global mean IRK peak sensitivity for August 2006 is about 0.6 mW/m<sup>2</sup>/ppb at around 550 hPa, decreasing linearly in pressure towards the surface and the tropopause. TES ozone and IRKs have been applied to chemistry climate model evaluation [Aghedo *et al.*, 2011a] and TES sampling has been shown to be sufficiently accurate to estimate zonal monthly mean distributions to within a few ppb [Aghedo *et al.*, 2011b]. The unweighted global mean of the all-sky longwave radiative effect, which includes both natural and anthropogenic ozone, is  $0.33 \pm 0.02$  W/m<sup>2</sup> [Worden *et al.*, 2011]. The longwave radiative effect from TES is less than most estimates of the anthropogenic component (i.e., radiative forcing) alone [Forster *et al.*, 2007]. These differences can be attributed in part to definitions of radiative forcing, which is commonly defined at the tropopause, spans the short-wave (SW) and long wave (LW) spectrums, and includes stratospheric temperature adjustment. The SW is generally a small contribution to ozone RF. The instantaneous TOA RF is about 10–20% higher than the stratospherically adjusted RF depending on the model [Forster *et al.*, 2007].

## 2.2. GEOS-Chem Forward and Adjoint Model

[5] GEOS-Chem ([www.geos-chem.org](http://www.geos-chem.org)) is a chemical transport model primarily driven by assimilated meteorology from the Goddard Earth Observing System (GEOS) of the NASA Global Modeling and Assimilation Office (GMAO). The spatial resolution of the GEOS meteorological fields are reduced to facilitate detailed simulation of tropospheric gas-phase HO<sub>x</sub>-NO<sub>x</sub>-VOC chemistry [Bey *et al.*, 2001]. For this work, we use model v8-02-01 with relevant updates through v9-01-01, run at the global  $2^\circ \times 2.5^\circ$  resolution. Global anthropogenic emissions of NO<sub>x</sub> are from EDGAR [Olivier *et al.*, 2001], overwritten by regional inventories in specific areas [van Donkelaar *et al.*, 2008]. Monthly biomass burning emissions are from GFEDv2 [van der Werf *et al.*, 2009] and biofuel emissions from Yevich and Logan [2003]. The adjoint of GEOS-Chem [Henze *et al.*, 2007] solves a set of equations auxiliary to the forward chemical transport model in a manner that efficiently yields the gradient of a scalar forward model response function with respect to all model parameters simultaneously. The adjoint has been used for analyzing long-rang impacts on O<sub>3</sub> [Zhang *et al.*, 2009; Walker *et al.*, 2012] as well as O<sub>3</sub> assimilation [Singh *et al.*, 2011; Parrington *et al.*, 2012].

## 2.3. Ozone Radiative Forcing

[6] We first define the mean area weighted observed outgoing TOA longwave radiative forcing:

$$\mathcal{J} = \frac{1}{A} \sum_i^N a_i F_i \quad (2)$$

where  $F_i$  is the  $i$ th of  $N$  TOA fluxes measured by the TES satellite. The product is weighted by the area of the model grid,  $a_i$ , and normalized by the total area,  $A = \sum_i^N a_i$ . The sensitivity of the mean TOA flux to emissions of each ozone precursor in each model grid cell is then

$$\lambda = \nabla_{\mathbf{E}} \mathcal{J} \quad (3)$$

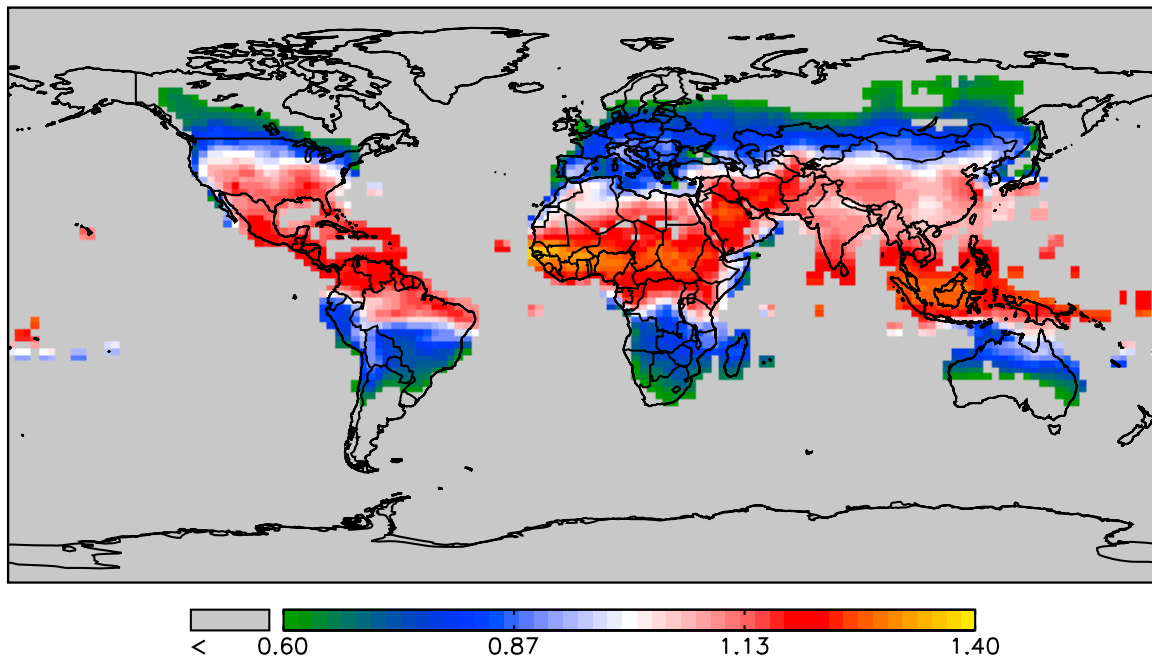
where  $\mathbf{E}$  is a vector of emissions from each species, sector and in each grid cell. Equation (3) is the direct ozone radiative forcing when  $\mathbf{E}$  is anthropogenic.  $\lambda_i$ , The sensitivity in equation (3) can be calculated for any single observed TOA flux from TES,  $F_i$  (extension to the global mean TOA flux,  $\mathcal{J}$ , is straightforward) as

$$\lambda_i = \frac{a_i}{A} \frac{\partial F_i}{\partial \mathbf{E}} = \frac{a_i}{A} \left( \frac{\partial \mathbf{c}_i}{\partial \mathbf{E}} \right)^T \frac{\partial F_i}{\partial \mathbf{c}_i}, \quad (4)$$

where the chain rule is employed to separate the sensitivity of flux to concentrations and the sensitivity of concentrations to emissions. The product of the two terms on the far right hand of equation (4) is calculated by the adjoint model [see Henze *et al.*, 2007]. The novelty of our approach is to use the observationally derived TES IRK's (equation (1)) to quantify the derivative of the observed flux with respect to retrieved ozone profiles, and to then propagate this sensitivity efficiently backwards in time using an adjoint model to obtain sensitivities with respect to emissions. The adjoint sensitivities of the August DRF are integrated backwards through the beginning of July, by which time they asymptotically approach steady state values owing to the lifetime of tropospheric O<sub>3</sub>. As shown in Figure S1, adjoint estimates for ozone DRF from the emissions in individual grid cells are in consistent agreement with evaluation of the full forward model across a wide range of emission perturbations with a slope of 0.991 and  $R^2 = 0.993$ , and the adjoint-based forcings are additive for modest changes to emissions across broader scales.

## 3. Results

[7] The sensitivities of the outgoing longwave radiation at the top-of-the-atmosphere (TOA) as observed by TES with respect to spatially-resolved anthropogenic NO<sub>x</sub>, CO, and



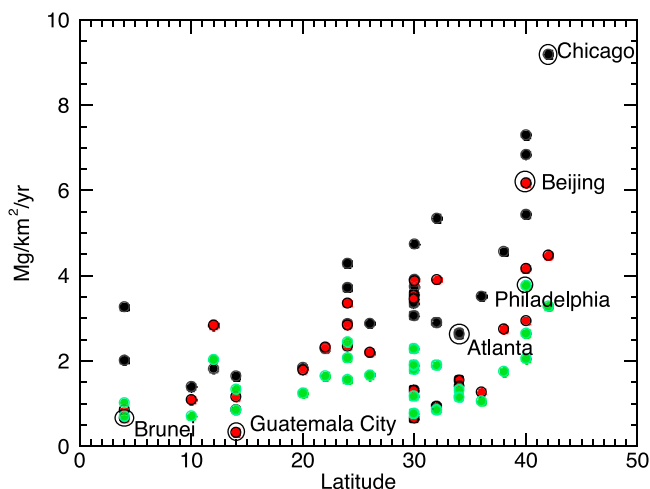
**Figure 2.** The impact of NO<sub>x</sub> emission locations on ozone DRF. Areas where the radiative effectiveness ratio,  $r_{eff}$ , is greater (less) than one indicate regions where additional NO<sub>x</sub> emissions would lead to an amplified (diminished) mean global ozone radiative forcing relative to the change in mean global ozone.

NMHC emissions are shown in Figure 1. These global emissions are defined on a  $2^\circ \times 2.5^\circ$  grid, which we refer to as “regions.” We note that this observationally derived radiative forcing at the TOA is significantly lower [Worden *et al.*, 2008] than typical values modeled at the tropopause [Forster *et al.*, 2007]. We have focused on August, 2006, which is the seasonal maximum in ozone radiative forcing for North America and consequently represents the strongest diversity of forcing responses to emissions [Naik *et al.*, 2005]. There are  $15 \times 2^\circ \times 2.5^\circ$  regions with a combined radiative forcing greater than  $0.15 \text{ mW/m}^2$ , which represents about 8% of ozone DRF from all anthropogenic emissions. For brevity, we denote these  $2^\circ \times 2.5^\circ$  regions by the names of the major city contained therein. Regions within China accounts for 10 of these and includes the Shanghai region, which is globally the most important:  $0.31$  ( $0.20$  (NO<sub>x</sub>),  $0.07$  (CO),  $0.04$  (NMHC))  $\text{mW/m}^2$ . While NO<sub>x</sub> emissions are usually the dominant radiative forcing in any single location, the Henan province centered near Zhoukou ( $34^\circ\text{N}$ ,  $115^\circ\text{E}$ ) is distinguished by a larger impact of CO and NMHC (56%) versus NO<sub>x</sub> emissions (44%) to its total radiative forcing of  $0.16 \text{ mW/m}^2$ . The US contribution includes Houston  $0.17$  ( $0.12$  (NO<sub>x</sub>),  $0.02$  (CO),  $0.03$  (NMHC)), New Orleans  $0.16$  ( $0.12$  (NO<sub>x</sub>),  $0.02$  (CO),  $0.02$  (NMHC)), and Western Atlanta  $0.16$  ( $0.11$  (NO<sub>x</sub>),  $0.02$  (CO),  $0.02$  (NMHC)). The remaining regions are Mexico City  $0.27$  ( $0.16$  (NO<sub>x</sub>),  $0.04$  (CO),  $0.07$  (NMHC)) and Kuala Lumpur  $0.15$  ( $0.13$  (NO<sub>x</sub>),  $0.01$  (CO),  $0.01$  (NMHC)). The impact of these latter regions is accentuated by their efficient transport pathways to the upper troposphere. Consistent with previous studies [Naik *et al.*, 2005], high latitude regions, such as in Europe, play a minor role in direct ozone radiative forcing despite having comparable emission levels.

[8] The heterogeneity in ozone DRF for August 2006 as shown in Figure 1 is a function of several factors: the season,

the magnitude of the underlying emissions in each grid-cell, the photochemical efficiency of O<sub>3</sub> formation in a given location per amount of precursor emitted, the transport of ozone into the free troposphere and the underlying distribution of clouds, water vapour, and temperature. In order to isolate the role of the physical atmospheric structure relative to the emission magnitude, we define a radiative forcing efficiency ( $r_{eff}(x, y)$ ) as a ratio of the global mean ozone DRF sensitivity for a single emission location to the global mean ozone sensitivity to that same emission location. If, for example, changing NO<sub>x</sub> emissions for a location by 100% leads to a 0.1% change in global mean ozone and a 0.125% change in global mean ozone DRF, then  $r_{eff} = 1.25$ .

[9] A plot of this radiative forcing efficiency is shown in Figure 2 for NO<sub>x</sub> emissions. Differences are particularly striking between North America and Europe, with NO<sub>x</sub> emissions from the former producing O<sub>3</sub> that is nearly twice as radiatively efficient. The meridional effect in radiative forcing efficiency [Naik *et al.*, 2005] leads to a maximum in the tropics, decreasing poleward by over a factor of two. In general, this ratio is highest in areas with convective lofting (i.e., the tropics) and over regions of higher altitude. However, there are important zonal variations that are related to cloud cover and convection. For example,  $r_{eff}$  is high ( $>1.2$ ) over Saudi Arabia and Iran for August 2006 because surface temperatures are both very high and the regions are relatively cloud free. Changes in atmospheric circulation due to monsoons have a significant impact on the export of surface emissions to the free troposphere. The highest values of  $r_{eff}$  are due to the Western African Monsoon, which is related to the shift of the inter-tropical convergence zone northward from the tropical Atlantic ocean towards the Saharan desert. The strong poleward temperature gradient results in a complex circulation pattern leading to significant convection and export along both mid-level African and high-level Tropical



**Figure 3.** NO<sub>x</sub> (black), CO (red), and NMHC (green) emissions that have about  $0.1 \pm 0.01$  mW/m<sup>2</sup> DRF. CO emissions have been reduced by 10 to fit on the same scale. Selected cities represent  $2^\circ \times 2.5^\circ$  metropolitan regions and are identified adjacent to their respective emissions.

Easterly Jets [Sauvage *et al.*, 2007]. The onset of the Indo-Australian Monsoon is influenced by El Niño conditions and a complex air-land interaction [Moron *et al.*, 2009], but the region is convectively unstable for all seasons leading to a very high  $r_{eff}$ . Radiative forcing efficiencies of CO and NMHCs, which are available in Figure S2, have the same meridional gradient but a much more diffuse zonal distribution.

[10] The variability of  $r_{eff}$  on spatial resolutions at which air quality strategies are enacted in practice has important policy implications. To illustrate the potential of this approach for supporting policy analysis,  $2^\circ \times 2.5^\circ$  regions with approximately the same total DRF, i.e., the DRF from the sum of NO<sub>x</sub>, CO, and NMHC emissions, of  $0.1 \pm 0.01$  mW/m<sup>2</sup> are shown in Figure 3 (details shown in Table S2 in Text S1).

[11] Across the 27 regions matching this criteria, NO<sub>x</sub> emissions represent about  $64 \pm 14\%$  of the DRF. Emissions increase with latitude by roughly a factor of 5–20 depending on the precursor emissions with considerable zonal scatter. For example, the Guatemala City region has 10 times smaller NO<sub>x</sub> emissions than Chicago and almost 20 times lower CO emissions than Beijing but correspond to the same total DRF. Similarly, the Brunei region near Malaysia has 5 times lower NMHC emissions than the Philadelphia region. These variations are driven by the poleward temperature gradient as well as cloud cover and large scale processes such as the Asian monsoon. Continental scale processes lead to a considerable spread in emissions as well. The region east of Atlanta (not to be confused with the Western Atlanta region discussed previously) has the lowest NO<sub>x</sub> emissions of the 11 selected regions and is 3.5 times lower than Chicago. The enhanced sensitivity in the Southeastern US is associated with summertime convection [Li *et al.*, 2005], as shown by  $r_{eff}$  in the Figure S3. Consequently, the variability of US emissions with  $0.1$  mW/m<sup>2</sup> DRF is about half of the mean:  $4.3 \pm 2.1$  (NO<sub>x</sub>),  $21 \pm 12$  (CO), and  $1.7 \pm 1.0$  (NMHC) Mg/km<sup>2</sup>/yr. The choice of metrics used in an air-quality climate co-benefit analysis could lead to very different results based on this variability. For example, a 10% reduction in

NO<sub>x</sub> emissions in Chicago would lead to  $0.01$  mW/m<sup>2</sup> change in DRF but the equivalent absolute reduction to emissions east of Atlanta would lead to a  $0.035$  mW/m<sup>2</sup> DRF reduction. Consequently, controlling against emissions for air quality versus DRF for climate can lead to very different strategies depending on location. It is important to identify various approaches to attaining the targeted O<sub>3</sub> DRF as reductions in NO<sub>x</sub> emissions will increase methane lifetime—a more efficient greenhouse gas—whereas reductions in CO and NMHC will decrease methane lifetime [West *et al.*, 2006].

[12] The accuracy of grid-scale radiative forcings is limited by a knowledge of precursor emission distributions and their ozone response, tracer transport, and the distribution of clouds. Nevertheless, such results can be aggregated and compared to previous studies. While the magnitude of the infrared, top-of-the-atmosphere ozone DRF reported here is smaller from ozone DRF defined at the tropopause by a factor of six, the relative sensitivity of ozone DRF to fractional emission changes aggregated to continental scales closely follows that of a previous work [Naik *et al.*, 2005] using different model emissions, chemistry and transport (see Figure S4). The agreement in relative sensitivity suggests that the differences are due to satellite versus model calculation of the ozone radiative effect rather than linearity assumptions in the adjoint approach.

#### 4. Conclusion

[13] Overall, we have shown here that there is substantial variability in the radiative forcing of O<sub>3</sub> precursor emissions at regional scales, and that the combined use of remote sensing observations and adjoint modeling provides a means of characterizing such variability. Further, there is considerable variability in the extent to which different precursors (NO<sub>x</sub> vs CO vs hydrocarbons) contribute to ozone's radiative impacts, as well as variability in the contribution of different emissions sectors within these species, which are critical for the overall O<sub>3</sub> response to emissions changes when accounting for the full range of chemical and physical feedbacks. Incorporation of climate co-benefits into air quality mitigation strategies thus requires quantitative understanding of chemical and physical processes at scales ranging from sub-continental to global.

[14] Equally important as design of control strategies is a framework to observe and assess the efficacy of these strategies against the backdrop of natural variability. While we do not address this framework explicitly, the satellite observations and assimilation system used in this study would be essential elements. The proposed suite of geo-stationary composition satellites from the Atmospheric Composition Constellation (ACC) as part of the Committee on Earth Observing Satellites (CEOS) (<http://www.ceos.org>) in conjunction with surface measurements could potentially provide the necessary observing system to support the implementation of ozone climate mitigation strategies. The requirements for such a system will be a point of future research.

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